

#### Contents lists available at ScienceDirect

## Journal of Affective Disorders

journal homepage: www.elsevier.com/locate/jad



## Review

# Acceptance and commitment therapy – Do we know enough? Cumulative and sequential meta-analyses of randomized controlled trials



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#### ARTICLE INFO

# Article history: Received 3 June 2015 Received in revised form 23 July 2015 Accepted 27 October 2015 Available online 30 October 2015

Keywords:
Sequential meta-analysis
Acceptance and commitment therapy
Mental health
Treatment efficacy
Anxiety
Depression

#### ABSTRACT

Acceptance and Commitment Therapy (ACT) has accrued a substantial evidence base. Recent systematic and meta-analytic reviews suggest that ACT is effective compared to control conditions. However, these reviews appraise the efficacy of ACT across a broad range of presenting problems, rather than addressing specific common mental health difficulties. Focussing on depression and anxiety we performed a meta-analysis of trials of ACT. We incorporated sequential meta-analysis (SMA) techniques to critically appraise the sufficiency of the existing evidence base. Findings suggest that ACT demonstrates at least moderate group and pre-post effects for symptom reductions for both anxiety and depression. However using SMA findings are more qualified. There is currently insufficient evidence to confidently conclude that ACT for anxiety is efficacious when compared to active control conditions or as primary treatment for anxiety. Similarly, using SMA, there is currently insufficient evidence to suggest a moderate efficacy of ACT for depression compared to active control conditions. To stimulate further research we offer specific estimates of additional numbers of participants required to reach sufficiency to help inform future studies. We also discuss the appropriate strategies for future research into ACT for anxiety given the current evidence suggests no differential efficacy of ACT in the treatment of anxiety compared to active control conditions

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#### 1. Introduction

Within the last decade third wave treatment approaches (Hayes, 2004a) have widened the spectrum of evidence-based psychological treatments, particularly in relation to mental health conditions deemed longstanding, complex or treatment resistant. Third wave' therapies have gained currency as an alternative to more established models of cognitive behavioural therapy (CBT) (e.g. Beck, 1963) via a relatively greater emphasis on context and experiential facets of psychological experience.

Third wave cognitive behavioural therapies include among others Dialectical Behavioural Therapy (DBT, Linehan, 1993 Mindfulness Based Cognitive Therapy (Segal et al., 2012), Compassion Focused Therapy (Gilbert, 2004), and Acceptance and Commitment Therapy (ACT, Hayes, et al., 1999). The third wave therapies also make explicit attempts to balance a coherent theoretical underpinning with a commitment to empirical testing.

Controlled trials have suggested efficacy for ACT in the treatment of depression, mixed depression and anxiety, physical health problems and psychotic disorders. Meta-analyses of randomized controlled trials of ACT have suggested a moderate to large effect size on primary outcomes measures after treatment and at followup (Hayes et al., 2006; Öst, 2008; Powers, Zum Vorde Sive Vording and Emmelkamp, 2009, Ruiz, 2010, 2012). A recent meta-analysis of ACT by Ruiz (2012) concluded that ACT outperformed CBT (Hedges g=0.4). However, the debate regarding the differential efficacy of ACT compared to other evidence-based psychological interventions is ongoing (e.g. Hofmann and Asmundson, 2008, 2010; Ost, 2009). The proliferation of third wave approaches raises questions for clinicians and policy makers (and clients/service users) regarding which therapeutic intervention is of optimal benefit for a given disorder or difficulty. This is especially important to the development of clear guidelines for the evidencebased practice of psychological interventions.

Existing evidence from systematic and meta-analytic reviews provide qualified support for the effectiveness of ACT as a psychological intervention when compared with no intervention (Ruiz, 2012; Powers et al., 2009). However, the data with regards to ACT in comparison to other psychological therapies are more equivocal. Therefore, clinicians, health service commissioners and policy makers at present must judge whether the evidence base for ACT is sufficient to make a confident recommendation regarding its efficacy. Borrowing from public health research (Muellerleile and Mullen, 2006; Wetterslev, Thorlund and Gluud, 2008), a novel statistical approach to this question is the appraisal of the sufficiency of the available cumulative knowledge. Where the total cumulative knowledge is still emerging, meta-analytic findings are at risk of false positives or false negatives due to methodological weaknesses such as power, random errors or systematic error (e.g. Kuppens et al., 2011). Sequential meta-analysis (SMA; Pogue and Yusuf, 1997) uses group sequential boundaries based on the alpha spending function to measure the accumulation of knowledge across studies, enabling decisions on the sufficiency of knowledge to recommend treatment to be made based on statistical properties. This approach, commonly used in the evaluation of medical interventions (e.g. Devereaux et al., 2005; Wetterslev et al., 2008) is under-utilised in the evaluation of psychological therapies. Although of potential benefit to evaluation of all evidence based psychological therapies we choose in this review to focus on ACT as an example of an emerging psychological therapy with a commitment to evidence-based practice.

In view of the above, our primary aim was to quantitatively review outcomes of ACT interventions for anxiety and depression using two complementary statistical approaches. Firstly, using cumulative meta-analytic techniques (CMA), we reviewed the evidence for ACT as a psychological intervention for anxiety and depression in group and pre-post comparisons. Secondly, we reviewed the evidence for the same conditions using sequential meta-analytic techniques (SMA). Use of SMA enabled us to make an estimate of the sufficiency of the evidence base for ACT. Secondary aims were to investigate the efficacy of ACT when compared against active treatments and when anxiety or depression were predetermined target outcomes. Regarding the primary aims we hypothesise that there is sufficient evidence to suggest that ACT is efficacious in the treatment of anxiety and depression. With regards to SMAs, to the best of our knowledge, this is the first time that a sequential meta-analytic approach has been used to appraise the sufficiency of evidence of ACT. Therefore no specific hypotheses were made.

#### 2. Method

Our quantitative review followed two stages. Firstly the literature was systematically searched to identify the study sample and to extract data. Secondly, the data was analysed using meta-analytic techniques. This stage incorporated conventional cumulative meta-analyses for ACT for anxiety or depression in group and pre-post comparisons, sequential meta-analyses for these conditions and lastly, subgroup analyses in which ACT was compared with active treatments and in conditions were anxiety or depression were predetermined treatment outcomes.

## 2.1. Literature search

## 2.1.1. Eligibility criteria

A systematic literature search was conducted to identify potential studies, following PRISMA guidelines (Moher, Schulz and Altman, 2008). Studies were included if they (1) investigated a manualised ACT approach, (2) used a randomised control design, (3) assessed anxiety or depressive symptoms using standardised outcome measures.

Studies were excluded if they (1) were not published in English, (2) did not include a standardised measure of anxiety or depression, (3) did not use an RCT methodology, or 4) were not published in a peer-reviewed publication, e.g. conference abstracts, book chapters, dissertations.

## 2.1.2. Information sources

Studies were identified by searching several database namely: PsycINFO 1840 to June 2015, MEDLINE 1966 to June 2015, SCOPUS 1841 to June 2013, Cochrane Central Register of Controlled Trials for 2014

#### 2.1.3. Search

We used search terms incorporating conjunctions of therapy and trial design terms: 'acceptance and commitment therapy' or 'acceptance', 'random', 'randomly', 'randomise', 'randomise', 'randomised', 'randomized', 'clinical trial', or 'trial'. These words were searched as key words, title, abstract, and MeSH subject heading terms. Also, we examined citation maps and used the 'cited by' search tools. Limits were then implemented to further refine the scope and ensure quality: databases were de-duplicated; searches limited to peer-reviewed articles; searches limited to human studies; and searches limited to adult studies. Reference lists of all relevant articles and existing systematic reviews (e.g. Powers et al., 2009) were screened by the authors to ensure no studies were overlooked.

#### 2.1.4. Data collection

The data was extracted into an electronic data extraction sheet by the first author and independently reviewed by the second author. Information extracted from the studies included (1) trial characteristics (including first author, publication year, and participant number), (2) control group characteristics (including active or non-active control), and (3) outcome characteristics (outcome measures).

## 2.1.5. Risk of bias measurement

To minimise the risk of data selection biases we included data pertaining to all reported anxiety and depression outcomes.

## 2.1.6. Summary measures

Standardised mean differences with heteroscedastic population variances (SMDH) for independent groups and for dependent groups were calculated (Bonett, 2008, 2009). Additionally, considering the small study samples Knapp and Hartung's adjustments (Knapp and Hartung, 2003) were calculated, yielding closer to nominal standard errors of the estimated parameters.

## 2.1.7. Study selection

Inclusion, exclusion criteria and search terms were specified a-priori.

The literature search identified initially 1865 studies. Study titles and abstract were screened against inclusion and exclusion criteria. Two independent reviewers conducted screening (first and second authors). Studies were assessed by considering the eligibility by the second author and revised by the first author. Disagreement was resolved through discussion.

In cases where insufficient data was reported in the primary study to apply meta-analytic methods we attempted to retrieve additional information from the first author of the published paper. Five authors were contacted to retrieve additional information. One author provided additional information. The review flowchart is illustrated in Fig. 1.

Quality assessments of the studies were evaluated using the 34-item Consort 2010 checklist (Moher et al., 2001). Each item is rated on a 3-point scale from 0 to 1, where 0=absent, 0.5=partial, 1=complete. Overall, Consort checklist scores range from 0 to 34. Higher overall scores suggest superior methodological rigour. The second and third author independently assessed a randomized subset of 10% of studies to check for inter-rater reliability. All studies were randomised using www.random.org list randomizer. Where there were ambiguities between the reviewers, the studies

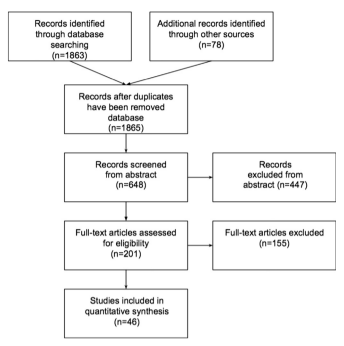


Fig. 1. Flow diagram of the study selection process.

were jointly reviewed to reach a unanimous decision. If further ambiguities remained a third reviewer had the ultimate decision. All remaining studies were randomised and divided amongst the two reviewers for independent evaluation. Overall the inter reviewer concordance was Kappa=0.85, suggesting an excellent agreement (Banerjee et al., 1999).

## 2.2. Literature appraisal

## 2.2.1. Meta-analytic approach

All meta-analytic procedures were conducted using the R-software package 'metafor' (Viechtbauer, 2015). Conducting consecutive meta-analyses that incorporate the same or virtually the same study sample can significant increase the risk of both type I and type II errors (Higgins et al., 2003; Borm and Donders, 2009). Findings may be erroneously deemed positive when they may be due to chance findings or false-positives. Similarly, findings might be rejected as negative or neutral due to lack of precisions or statistical power. A sequential meta-analysis method offers a statistical approach to manage these risks (Wald, 1947; Pogue and Yusuf, 1997).

In this analysis we used a four-steps analytic approach described by Kuppens and Onghena (2012) including the calculation of the optimal information size, conducting cumulative meta-analyses, constructing sequential boundaries, and determining statistical sufficiency. These steps will be described in detail in the ensuing sections.

2.2.1.1. Computing optimal information size (OIS). Arguably the level of convincing evidence for a meta-analysis should be no less than that of a well-designed single trial. To determine the threshold level of sufficiency Pogue and Yusuf (1997) proposed that conventional sample size calculation methods can be used. The optimal information size (OIS) is the sample size required to detect a presumed pooled effect whilst minimising type I and II errors. However, this approach does not control for heterogeneity between studies when using meta-analytic methods. Therefore a heterogeneity adjusted method for hierarchical designs can be used to control for the degree of between study variability

(Higgins et al., 2010; Kuppens and Onghena, 2012). The heterogeneity adjusted OIS (HOIS) can be readily calculated as follows:

$$HOIS = \frac{OIS}{(1 - I^2)}$$

Conventionally, the  $I^2$  index measures the extent of true heterogeneity dividing the difference between the Q value and its degrees of freedom (k-1) by the Q value, and multiplied by 100 (Borenstein, 2009). To render authorities conclusions regarding the evidence base for ACT we chose more stringent criteria for alpha, power and proposed effect sizes ( $\alpha$ =.01,  $1-\beta$ =.9) as recommended by Kuppens and Onghena (2012). Considering the moderate to high heterogeneity reported in previous meta-analyses (Ruiz, 2012) the authors also elected to use a high heterogeneity estimate (e.g.  $I^2$ =75%) in their HOIS calculations. Lastly, considering currently reported effect size estimates a medium effect size (e.g. d=.5) was selected to be a reasonable estimate for the HOIS calculations. We used the open-source programme G\*Power 3 (Faul et al., 2007) to calculate heterogeneity adjusted OIS's (Table 3).

2.2.1.2. Cumulative meta-analysis. To obtain pooled cumulative estimates of effect sizes over time cumulative meta-analyses were conducted on a chronologically ordered study sample (e.g. publication year). Random models were used to calculate pool effect sizes at interim analysis points.

Independent meta-analyses were conducted for anxiety and depression in order to conform to the assumption of independent effect sizes that underline meta-analytic procedures (Borenstein, 2009). When several relevant outcome measures were reported, the outcomes yielding the most conservative effect size estimate per study were included in the analyses.

2.2.1.3. Sequential boundaries. The optimal information size can be used to calculate the group sequential boundaries b using the alpha-spending function by Lan and DeMets (1983). This function  $\alpha^*$  is a monotonically non-decreasing function which allocated the allowable Type I error through a function based on the information fraction t. The information fraction t in turn is the proportion of (heterogeneity adjusted) optimal information size (HOIS) that has been accumulate at a particular interim analysis point q thus  $t_q = \frac{i_q}{HOIS}$ . Several functions can be fitted into the Lan and DeMets (1983) alpha spending function including the O'Brien Fleming used in this analysis (Reboussin et al., 2000):

 $\Phi$  is the standard nomal distribution function. The function thus allows for the calculation of sequential boundaries  $b_q$ . This statistic corresponds to the critical *Z*-value for the allocated  $\alpha$  at each step of the interim analysis.

2.2.1.4. Sufficiency. At each interim point of the analysis q we obtain two standardized test statistics Zq and  $b_q$  that denote the Z values of the pooled effect size and allocated  $\alpha$  respectively. Sufficiency is determined by comparing the Zq and  $b_q$ scores at each step of the interim analysis. As long as  $[Zq] < b_q$  sufficiency has not yet been attained and further studies are needed. However, if the criterion of  $[Zq] \ge b_q$  is reached at an interim point of the analysis sufficiency of evidence indicates that a predetermined treatment effect exists. In turn, if the optimal information size has been reached and  $[Zq] < b_q$ , sufficiency of cumulative evidence is achieved to refute the effectiveness of the intervention in relation to the predetermined effect size,  $\alpha$  and power.

#### 3. Results

#### 3.1. Literature search

The systematic literature search identified k=28 and k=39 eligible randomized controlled trials studies of ACT for anxiety and depression respectively. Study characteristics including trial characteristics, control group characteristics, and outcomes are described in Tables 1–3.

The total participant sample in this quantitative review was n=1628 and n=1987 participants in anxiety and depression trials. The sample size for participants within both anxiety and depression treatment trials varied from n=6 to n=125 per group. All papers were published between 1989 and 2015. Within the reviewed sample the modus of published trials for anxiety treatment were in 2011, 2013 and 2014 (k=8). In turn, for depression the modus of published depression treatment trials was in 2011 and 2012 (k=8). For both anxiety and depression outcomes the majority of trials compared ACT against a waiting list control (WL; k=17 and k=22 studies for anxiety and depression respectively). The majority of anxiety treatment trials used CBT as bona fide treatment comparator (k=7). Similarly, the most frequent bona fide control treatment condition in depression treatment trials was CBT (k=9). The most frequently used anxiety outcome measure was the Depression Anxiety and Stress Scale (DASS-A, k=9). In turn, the most frequently used depression outcome measure was the Beck Depression Inventory (BDI, k=18). Less than a third of anxiety trials (k=10) employed an active control condition compared to 38% of depression trials (k=15). Eighteen per cent of anxiety trials (k=5) and 31% of depression trials (k=12) predetermined the primary target outcome.

## 3.2. Literature appraisal

#### 3.2.1. Optimal information size (OIS)

The calculated heterogeneity adjusted optimal information size was n=220 and n=848 for pre-post and group comparisons respectively.

## 3.2.2. Cumulative meta-analyses

A series of random-effects cumulative meta-analyses were conducted. Findings are listed in Table 4. Regarding the primary aims the cumulative meta-analysis yielded large significant effects for pre-post treatment reduction in anxiety (d=.95, p < .001) and depression (d=.92, p < .001) scores. Analyses also revealed a small significant effect for group treatment changes for anxiety (d=.45, p < .05) and a medium effect size for depression (d=.54, p < .001) favouring ACT.

Regarding the secondary aims, for ACT for anxiety the pre-post comparison suggested a large significant effect (d=1.85, p<.01) when anxiety was predetermined as primary treatment target. In turn, ACT for anxiety in active control conditions revealed no effect (d=-.04, n.s.). Similarly, findings for ACT for anxiety in group comparisons when anxiety was the primary treatment target revealed a non-significant large effect (d=-.77, n.s.). In pre-post comparisons for ACT where depression was the primary treatment target findings suggested a large significant effect (d=1.22, p<.001). Group comparisons for ACT for depression as a primary treatment target suggested also large significant effects (d=.73, p<.001). In group comparisons for ACT versus an active control conditions findings suggested a small non significant effect (d=.26, n.s.)

#### 3.2.3. Sequential meta-analyses

In relation to sufficiency, findings will initially be described for ACT in both pre-post and group comparisons for the whole study sample. We will then continue to describe findings comparing ACT

 Table 1

 Characteristics of studies included in the meta-analyses.

Study reference	Country and type of sample	Recruitment	Initial sample size (ana- lysed sample)	Mean age (s.d)	Male/ female (n)	Ethnicity %	Education	Problem and severity	Treatment Dosage for ACT intervention
<b>Studies Measuring</b> Zettle and Rains (1989)	Depression as outco USA Community	<b>Dime of interest</b> Self-selecting Volunteers via media	31 (31)	41.3 (N/R)	0/31	N/R	Mean 14.1 yearsof education; All at least High School Level	Clinical Depression	12 weekly group sessions
Bond and Bunce. (2000)	UK Business	Self-selecting volunteers via flyers	90 (65)	36.43 (9.72)	45/45	N/R	Primarily University Graduates	Stress Management	3 group sessions over 14 weeks
Hayes et al. (2004b)	USA – Clinical: Substance Misuse in Treatment	Recruitment from 3 Methadone Clinics	138 (78)	42.2 (N/ R)	67/72	Ethnic Minorities = 13%	N/R	Participants met DSM-IV cri- teria for Substance Abuse or Dependence	16 week protocol, 48 sessions  – 32 individual +16 group
Lappalain et al. (2007)	Finland - clients seeking psychotherapy	Self-selecting from media adverts	28 (28)	41.8 (13.2)	3/25	N/R	N/R	Common difficulties pre- dominatly depression, inter- personal and anxiety problems	10 individual sessions
Wicksell et al. (2009)	Sweden - Clinical sample of Adoles- cents with chronic pain	Chronic idiopathic pain referred to specialist Pain Treatment Service	32 (32)	14.8 (2.4)	7/25	N/R	N/R	Chronic pain of > 3 months	10x weekly 1hour individual sessions + 1 – 2 sessions with parents (90 mins)
Hinton et al. (2010)	USA – University Students	Self selecting volunteers via flyers	22 (22)	20.09 (2.56)	6/16	Euro/American = 86.5% African/American = 4.5% MultiEthnic = 9%	96% full time college students	Evidence of low-self esteem or negativity AND scoring as Distressed on standardised measures	3 weekly, 1 hour individual sessions
Smout et al. (2010)	Australia -Clinical sample of Metham- phetamine Users	Recruited from Drug and Alcohol Services, plus media recruitment	104 (31)	30.9 (6.5)	57/47	N/R	7-10 years =25% 11-13 years =49% Vocational educa- tion =17% University =9%	Met DSM-IV criteria for me- thamphetamine abuse or de- pendence; Use at least once weekly over past 3 months	12x weekly 1 hour individual sessions
Twohig et al. (2010)	USA – Community sample of in- dividuals present- ing with OCD	Recruitment from Health Professionals and via media	79 (79)	37 (15.5)	31/48	Caucasian =88.6% African American = 1% Asian American = 2.5% Latin American = 5% Native American = 2.5%		Met DSM-IV criteria for OCD	8x weekly 1 hour individual sessions
Hayes et al. (2011)	Australia – Clinical Sample of adolescents	Recruitment from public child and adolescent psychiatric services	38 (30)	14.9 (2.55)	11/27	N/R	71% of sample at- tending school	Moderate to severe depressive symptoms via DAWBA	Individual sessions (no number of sessions given)
Folke et al. (2012)	Sweden – In- dividuals on dis- ability or illness benefits	Recruitment from re- gional Social Insurance Office	34 (27)	43.24 (9.46)	4/30	All Caucasian	N/R	DSM-IV Unipolar Depression+Unemployment and Sick Leave	1 individual+5 group sessions
Kocovski et al. (2013)	Canada – Commu- nity sample of So- cial Anxiety Disorder	Recruitment from Health Professionals and via media	137 (137)	Tx: 34.94 (12.52)	63/74	White = 62% Asian = 20% Black = 3.6% Hispanic = 3.6% Other = 10.9%	College or university=63.5% Some post secondary education=27.0% N/R=9.5%	DSM-IV diagnosis of Social Anxiety Disorder, Generalised	12x weekly 2 hour individual sessions
Alonso et al. (2013)	Spain – selected clinical sample in nursing care homes	Recruitment from two selected nursing homes	10 (10)	Tx: 87 (2.44)	2/8	N/R	None=40% Primary=50% Secondary=20%	Chronic musculoskeletal pain of articular origin for $> 6$ months	10x twice weekly group sessions of 2 hours per week.
Lappalainen et al. (2013)	Finland – commu- nity sample of males	Self-selecting via news- paper advert	24 (24)	Tx: 47.1 years (SD 4.72	24/0	N/R	Tx: Mean duration of 7.1 years	"Exhaustion, stress symptoms, or sleeping problems".	Integrated programme of web/mobile apps, personal monitoring and software

Study reference	Country and type of sample	Recruitment	Initial sample size (ana- lysed sample)	Mean age (s.d)	Male/ female (n)	Ethnicity %	Education	Problem and severity	Treatment Dosage for ACT intervention
									with 3x group intervention
McCracken et al. (2013)	UK – Community sample with Chronic Pain	Recruitment from General Medical Practices	73 (58)	58.0 (12.8)	23/50	White British=97.3%		Persistent pain of longer than 3 months' duration with GP consulations, Distress/Dis- ability and use of Medication	sessions 4x Group sessions, each4 hours long. 3x one week and a further session one week later.
Clarke et al. (2014)	UK-clinical sample of individuals with treatment resistant mental health problems	Referrals from Specialist personality disorder clin- ic in a public health setting	45 (45)	43.46 (s. d. 1/4 12.35).	20/41	N/R	N/R	Treatment resistance via > one previous 8-session episode of psycholo- gical therapy	16x weekly group sessions of 2 hours duration
Livheim et al. (2014)	Australia – com- munity sample of young people at school	Referral via school counsellors	51 (51)	14.6 (1.03)	8/43	N/R	N/A	mild to moderate depressive symptoms	8x weekly group intervention sessions
Tamannaeifar et al. (2014)	Iran – Clinical sam- ple of women with depressive disorder	Referral to University Clinic	19 (19)	25.2 (4.2)	0/19	N/R	Diploma=58% Master=42%	DSM-IV diagnosis of Major Depressive Disorder	12x twice weekly group intervention sessions
	Iran -clinical sam- ple of women with breast cancer	Referrals from specialist clinic	30 (30)	N/R	0/30	N/R	N/R	Depressive symptoms, no diagnostic or severity criteria	8 intervention sessions of one hour duration
Kohtala et al. (2015)	Finland - commu- nity sample of in- dividuals reporting depressive symptoms	Self-selecting via local media	57 (57)	46.2 (SD=11.9	12/45	All caucasian	Comprehensive school=9% Secondary school=45% Higher education=43% Other=3%	subjective depressive symptoms or depressed mood	4x group intervention sessions of 1 hour duration
	<b>Anxiety as outcome</b> USA-College	Self selecting volunteers	37 (24)	30.9 (N/	7/30	White= 66.6%	N/R	Test anxiety	6 individual weekly sessions
Brown et al. (2011)	students USA-University students	via flyers Self selecting volunteers from Psychology courses	16 (16)	R) 20.2 (1.9)	5/11	Black=21%Hispanic =12.5% White =43.7% Asian/Pacific Islander =25% Black =6.2% Caribbean/Haitian = 6.2% Latino=6.2% Multiracial/other=12.5%	N/R	Test anxiety	Single – 2 hour group session
Mo'tamedi et al. (2012)	Iran – clinical sam- ple of female pa- tients with headache	Self-referral from women attending specialised headache clinic	30 (30)	Tx: 34.18 (7.30)	0/30	N/R	Tx Group: 13.00 (2.90)	International Classification of Headache Disorders diag- nosis of primary chronic headache	8x weekly group sessions
Arch et al. (2012)		Adult Outpatients recruited via media	128 (128)	37.93 (11.70)	61/67	White =67.2% Asian American/ Pacific Islander =8.0% African American/Black =8.8% Hispanic/ Latino=12.0% American Indian/ Alaskan Native=0.08%	15.41 (2.07)	DSM-IV Diagnosis of Anxiety Disorder (Panic, Social Anxi- ety, Specific Phobia, OCD or GAD)	12 individual weekly 1-hour sessions
Zargar et al. (2012)	Iran-Clinical Sam- ple of GAD	Patients receiving treatment for GAD	24 (18)	Tx: 34.5 (2.41)	0/24	N/R	Guidance School = 11.1% High School = 77.8% Bachelor = 11.1%	DSM-IV Diagnosis of Generalised Anxiety Disorder	12 × 90 minute sessions
Craske et al. (2014)	USA – individuals with social phobia	Referrals from local fly- ers, Internet and local	87/87	28.37 (6.76)	47/87	White = 50.57% Hispanic/Latino = 17.24%	Mean no. of years = 15.04 (1.95)	DSM-IV Diagnosis of Social Anxiety Disorder	12 weekly sessions of in- dividual intervention

		newspaper advertise- ments, and referrals.				African American/Black=2.30% Asian American/Pacific Islander=18.39%			
Lanza et al. (2014)	population of wo- men with sub- stance misuse disorders	Referrals from Prison Team	50 (50)	Tx: 31.1 (6.4)	0/31	N/R	N/R	DSM-IV diagnosis of substance misuse disorder	16x weekly group sessions of 90 minutes duration
Studies measuring Gratz and Gunder- son (2006)		Anxiety as outcomes Patients referred from clinicians at psychiatric hospital, private practice and self-referrals via advert	22 (22)	33.32 (9.98)	0/22	All White	Some college =21% College Gradu- ate42% Graduate school=37%	Meeting 5 or more criteria for DSM-IV BPD+at least 1 epi- sode of DSH over previous 6 months	14x weekly group session
Woods et al. (2006)	USA – Unclear sampling	N/R	28 (25)	35.0 (10.2)	3/25	Caucasian = 96.4% African American = 3.6%	Mean years of education = 15.0 (2.8)	DSM-IV diagnostic criteria for Trichotillomania;	10 x individual sessions across 12 weeks
Forman et al. (2007)	USA – Clinical sam- ple with "Distres- sing Symptoms"	Clients presenting to University Counselling Centre	101 (99)	27.87 (7.25)	20/81	Caucasian =64.4% Asian =10.9% Black =12.9% Latino=3.0%	N/R	DSM-IV diagnostic criteria for depressive, anxiety or ad- justment disorders	Mean of 15.27 individual sessions
Roemer et al. (2008)	USA – clinical sam- ple with General- ised Anxiety Disorder	Clients seeking treatment at specialist centre for anxiety disorders	31 (31)	33.59 (11.74)	9/22	White=87% Latino/Latina=6.5 Black=3.25% Asian=3.25%	N/R	DSM-IV Diagnosis of Generalised Anxiety Disorder	$4 \times 90$ minute individual sessions $+ 12x$ individual weekly 1-hour sessions
Wicksell et al. (2008)	Sweden – Clinical sample of chronic pain and whiplash	Self-selecting via Patient Organisation in one geo- graphical area of Sweden	22 (20)	Tx: 48.2 (7.8)	5/16	N/R	N/R	Pain duration of > 3 months	10 individual sessions over 8 weeks
Johnston et al. (2010)	New Zealand – Clinical sample with chronic pain	Referral via contact with Clinical Psychology and Pain Clinics	24 (14)	Median age=43	10/14	N/R	N/R	No severity criteria	Self-Help intervention via book and workbook
Bohlmeijer et al. (2011)	Netherlands - Clin- ical sample with depressive symptoms	Self-referral via targeted recruitment from mental health institutions	93(93)	49.02 (10.70)	17/76	White Dutch=85% Other=8% N/R=7%	< 13 Years of education= 26.9% 13-16 Years of education=33.3% > 16 Years of education=39.8%	Mild to moderate psychological distress	$8 \times 2$ -hour group based intervention.
Fledderus et al. (2012)	Netherlands – Community sample with depressive symptoms	Self-referral via media adverts	376 (376)	42 (N/R)	113/ 263	White Dutch=93% Other=7%	Low Level=1.5% Middle Level =12% High Level=86.5%	Mild to moderate depressive symptoms	9-week protocol with Self- help book+weekly email support with guided questioning
Muto et al. (2011)	USA – Japanese students studying at overseas University	Self-referral via flyers on campus and email to students	70	23.6 (N/ R)	26/44	N/R	N/R	No severity criterion	Self-help workbook completed over 8 weeks.
Thorsel et al. (2011)	Sweden - clinical sample of in- dividuals with chronic pain	Recruitment from spe- cialised pain clinic	90(90)	46.0 years (SD 12.3)	32/58	N/R	N/R	Chronic pain with no severity criterion	1x initial individual face to face session; workbook+7 weeks of 30 minute phone support; 1 × 90 minute concluding face to face session
Westin et al. (2011)	Sweden – Clinical sample of in- dividuals with tinnitus	Recruited from audiology departments and self-re- ferral via adverts	64 (62)	50.9 years (SD=12.9	34/30	N/R	N/R	Tinnitus of duration of $> 6$ months	Up to 10x individual sessions, mean number of session=8.38 (1.56)
Wetherell et al. (2011)	USA – community sample with chron- ic pain	Recruited via clinics, advertisements, media, pain support groups, other studies, referrals from other participants	114 (114)	54.9 (12.5)	56/58	N/R	44.7% had at least a bachelor's degree	Chronic non-malignant pain > 6 months duration, with significant severity and interference	8x weekly group sessions
Jeffcoat and Hayes (2012)		Self-referral after mail drop via staff support	236 (186)	N/R	N/R	N/R	N/R	No severity criterion	Self-help workbook completed over 8 weeks.

Study reference	Country and type of sample	Recruitment	Initial sample size (ana- lysed sample)	Mean age (s.d)	Male/ female (n)	Ethnicity %	Education	Problem and severity	Treatment Dosage for ACT intervention
Jensen et al. (2012)	educational district Sweden – clinical sample of women with Fibromyalgia	office Referral via Primary Care Physicians	43 (34)	45.6 years (SD 6.4)	0/43	N/R	N/R	Meeting 1990 American Col- lege of Rheumatology diag- nostic criteria for fibromyalgia	12 weekly sessions (90 minutes each), conducted in groups of 6 patients.
Morton et al. (2012)	Australia – clinical sample of in- dividuals with Bor- derline PD	Referrals from specialist public sector mental health service	41 (28)	Tx: 35.6 (9.33)	3/38	N/R	Did not complete high school=29% Completed high school=34%Some tertiary/ has degree=37%		$12 \times 2$ -hour group intervention sessions
Buhrman et al. (2013)	Sweden - clinical sample of in- dividuals with chronic pain	Recruitment from spe- cialised pain clinic	76 (76)	49.1 (10.34)	31/45	N/R	Nine-year compul- sory school=9.2% Upper secondary school=47.4% University education= 43.4%	Functional impairment caused by chronic pain with medical investigation in previous year.	7 section online treatment programme with down- loadable mp3 files. Mean number of 4.2 (2.7) sections completed
Carlbring et al. (2013)	Sweden – commu- nity sample of in- dividuals reporting as depressed	Recruitment via news- paper advert	80 (80)	44.4 (13.5)	14/66	N/R		Symptoms of mild to moderate depression on MADRS-S	Module internet-based treatment programme + 15 minutes per week acces to internet therapist
Levin et al. (2014)	USA – Under- graduate college students	Self-selecting via campus and local adverts	76(76)	18.37 (.54)	35/41	Caucasian =71.1% African American =7.9% Asian =9.2% Latino/Hispanic =15.8% Native American=9.2% Pacific Islander/Hawaiian=2.6%	N/R	No severity criterion	2x web based multimedia lessons with tailored emails
Avdagic et al., 2014	Australia – in- dividuals present- ing with GAD	Self-selecting via campus and local adverts	51 (51)	36.17 (13.1)	17/34	N/R	N/R	DSM-IV Diagnosis of Generalised Anxiety Disorder	6x weekly group sessions of 2 hours duration
Livheim et al. (2014)	Sweden -commu- nity sample of young people at school	Self-selecting sample scoring above the 80th percentile SDQ, PSS and GHQ-12 at screening.	32 (32)	14–15 years	9/22	N/R	N/A	As for recruitmenbut not severe problems or suicidal ideation	$6 \times 90$ minute weekly group intervention sessions
Yadavaia et al. (2014)	USA-Under- graduate college students	Self-selecting sample of psychology under- graduates meeting screening criteria	78 (73)	19.69 (2.660	19/54	Asian/Pacific Islander = 16% African-American/Black = 7% Hispanic/Latino = 12% Native American = 1% White/Non-Hispanic = 74%	N/R	GHQ-10 score of < 10.	1 × 6 hour workshop

Notes: N/R=Not Reported; N/A=Not Applicable; Tx=Treatment Group; DAWBA=Development and Wellbeing Assessment (Goodman et al. 2000); UK=United Kingdom; USA=United States of America; GAD=Generalised Anxiety Disorder; OCD=Obsessive Compulsive Disorder; SDQ=Strengths and Difficulties Questionnaire; PSS=Perceived Stress Scale; GHQ-12=General Health Questionnaire – 12 item;

**Table 2**Measurement characteristics of ACT studies included in the meta-analysis for anxiety.

Study reference	Post treatme	ent group comparison	Outcome measure	Control condition	Control group	Outcome category	
	N (Tx) N (Control)		•				
Zettle (2003)	12	12	STAI	Active	SysD	Primary	
Gratz and Gunderson (2006)	12	10	DASS-A	Passive	WL	No Differentiation	
Woodset al. (2006)	12	13	PAI-A	Passive	WL	No Differentiation	
Forman et al. (2007)	55	44	BAI	Active	CT	No Differentiation	
Roemer et al. (2008)	15	16	DASS-A	Passive	WL	Primary	
Wicksell et al. (2008)	11	10	HADS-A	Passive	WL	Secondary	
Johnston et al. (2010)	6	8	BAI	Passive	WL	No Differentiation	
Bohlmeijer et al. (2011)	49	44	HADS-A	Passive	WL	Secondary	
Brown et al. (2011)	6	6	STAI	Active	CT	No Differentiation	
Fledderus et al. (2012)	125	126	HADS-A	Passive	WL	Secondary	
Muto et al. (2011)	30	31	DASS-A	Passive	WL	Secondary	
Thorsell et al. (2011)	28	27	HADS-A	Active	AR	No Differentiation	
Westin et al. (2011)	21	21	HADS-A	Active	TRT	Secondary	
Wetherell et al. (2011)	49	50	PASS	Active	CBT	No Differentiation	
Arch et al. (2012)	71	57	PSWO	Active	CBT	Primary	
Jeffcoat and Hayes (2012)	39	42	DASS-A	Passive	WL	Secondary	
Jensen et al. (2012)	19	15	STAI	Passive	WL	Secondary	
Mo'tamedi et al. (2012)	11	15	STAI-T	Passive	WL	No Differentiation	
Morton et al. (2012)	14	14	DASS-A	Passive	WL	Secondary	
Zargar et al. (2012)	9	9	PSWO	Passive	WL	No Differentiation	
Buhrman et al. (2013)	29	32	HADS-A	Passive	WL	Secondary	
Carlbring et al. (2013)	40	38	BAI	Passive	WL	No Differentiation	
Craske et al. (2014)	33	29	CSR	Active	CBT	Primary	
Lanza et al. (2014)	18	19	ASI	Active	CBT	No Differentiation	
Levin et al. (2014)	37	39	DASS-A	Passive	WL	No Differentiation	
Avdagic et al. (2014)	19	19	DASS-A	Active	CBT	Secondary	
Livheim et al. (2014) Swedish Sample.	15	15	DASS-A	Passive	TAU	Primary	
Yadavaia et al. (2014)	43	39	DASS-A	Passive	WL	No Differentiation	

Note. Outcome measure abbreviations: STAI – State Trait Anxiety Inventory, DASS – Depression Anxiety Stress Scale, PAI-A-Personality Assessment Inventory Anxiety, BAI – Beck Anxiety Inventory, HADS – Hospital Anxiety and Depression Scale, PASS – Pain Anxiety Symptom Scale, PSWQ – Penn State Worry Scale, AS-Control condition abbreviations: SysD – Systematic Desensitization, WL – Waiting List, CT – Cognitive Therapy, AR – Applied Relaxation, TRT – Tinitus Retraining Therapy, CSR=Clinical Severity Rating; ASI=Anxiety Sensitivity Inventory.

when anxiety or depression was specified a-priori as target outcome and against active control conditions. Lastly, findings from auxiliary analyses will be described.

As illustrated in (Fig. 2a, c and d), sufficiency was reached for all SMAs for anxiety and depression with the exception of ACT for anxiety in group comparisons (Fig. 2b). In this comparison, although statistical sufficiency (HOIS) was reached, the threshold boundary was not crossed thus based on these data it cannot be assumed that ACT is moderately effective in treatment of anxiety in group comparisons. It is of note that for ACT for depression in pre-post comparisons, the intermittent Z-value crossed the threshold boundary at some point in the analysis (Fig. 2c). However, at the end point when the accumulated alpha has been reached (i.e.  $t \ge 1$ ), the sequential boundary was not crossed ( $Z_{\alpha=.01}=2.46$ ,  $Z_{t\ge -1}=2.17$ ). Although at the end point of the analysis the sequential boundary has not been crossed, nonetheless statistical criterion for ACT as an efficacious intervention has been met.

Our analyses suggest that ACT for anxiety as an a-priori comparator in pre-post comparisons is at least moderately effective as the threshold boundary was crossed at some point in the analysis (Fig. 3a). In contrast, findings from the SMAs of ACT for anxiety in group comparisons when a-priori determined as treatment or when compared with active control conditions, suggest that there is currently insufficient evidence to indicate a medium effect (Fig. 3b and c). As described in Table 3 additional samples of n=573 and n=290 participants respectively would be required to reach the predetermined heterogeneity adjusted optimal information size for group comparisons when anxiety has been predetermined or compared to an active control condition.

In relation to ACT for depression findings suggest that there is sufficient evidence of a least a medium effect in studies where depression was a-priori specified as a target outcome. This is the case for both pre-post and group comparisons (Fig. 4a and b). Conversely, our findings suggest that ACT for depression in group comparisons with active control conditions there is currently insufficient evidence to indicate a medium effect as sufficiency (HOIS) has been reached (HOIS) (Fig. 4c). An additional samples of n=93 would be required to reach the predetermined heterogeneity adjusted optimal information size for group comparisons with active controls in depression.

## 3.2.4. Auxiliary analyses

To further explore the differential efficacy of ACT and CBT we conducted additional post-hoc meta-analyses. For anxiety findings from a cumulative meta analysis (k=8) suggest that there is currently no differential effect between these two treatment modalities (d=.08, n.s.). An additional n=260 participants would be required to be able to determine the sufficiency of cumulative evidence.

Similarly the cumulative evidence (k=10) of ACT for depression revealed no differential effect between ACT and CBT (d=-.01, n. s.). An additional n=314 participants would be required to be able to confidently appraise the statistical sufficiency of the cumulative evidence.

## 3.2.5. Publication Bias Analysis

As we did not include unpublished work in our meta-analyses we tested for publication bias in the following ways. Firstly, funnel plots were examined. demonstrating no systematic publication biases (tables available from first author on request). Secondly.Failsafe N (Rosenthal, 1979) values ranged from k=2048 to k=6708 studies for the primary study aims, suggesting absence of publication bias. For the secondary study aims Fail-safe N values

Table 3 Measurement characteristics of ACT studies included in the meta-analysis for depression.

N (Tx)  11 24 42 12	N (Control)  10 21 42	BDI			
24 42	21			_	
42			Active	CT	Primary
	42	BDI	Active	IPP	No Differentiation
12	42	BDI	Active	ITSF	Secondary
	10	DASS-D	Passive	WL	No Differentiation
12	12	PAI-D	Passive	WL	No Differentiation
55	44	BDI	Active	CT	No Differentiation
14	14	BDI	Active	CBT	Secondary
15	16	BDI	Passive	WL	Secondary
11	11	HADS-D	Passive	WL	Secondary
15	14	CES-DC	Active	MDT	Secondary
10	12	BDI	Passive	WL	Primary
6	8	CMDI	Passive	WL	No Differentiation
14	17	BDI	Active	CBT	Secondary
36	32	BDI	Active	PRT	Secondary
49	44	CES-D	Passive	WI.	Primary
125	126		Passive	WI.	Primary
	11		Passive	WL	No Differentiation
30	31	DASS-D	Passive	WL	Secondary
28	29	HADS-D	Active	AR	No Differentiation
21	18	HADS-D	Active	TRT	Secondary
					No Differentiation
					Primary
					Secondary
					No Differentiation
					Tertiary
					Primary
		•			Secondary
					Primary
					No Differentiation
					Secondary
					No Differentiation
					Primary
	14 15 11 15 10 6 14 36 49 125 19	14     14       15     16       11     11       15     14       10     12       6     8       14     17       36     32       49     44       125     126       19     11       30     31       28     29       21     18       49     50       14     13       45     44       20     16       14     14       29     32       40     38       37     32       31     27       5     5       11     12       24     15       32     19       15     17       10     9       37     39       19     19       28     39	14       14       BDI         15       16       BDI         11       11       HADS-D         15       14       CES-DC         10       12       BDI         6       8       CMDI         14       17       BDI         36       32       BDI         49       44       CES-D         125       126       CES-D         19       11       RADS-2         30       31       DASS-D         28       29       HADS-D         21       18       HADS-D         49       50       BDI         14       13       BDI         45       44       DASS-D         20       16       BDI         14       14       DASS-D         29       32       HADS-D         40       38       BDI         37       32       BDI         31       27       PHQ-9         5       5       GDS-10         11       12       BDI         32       19       RADS-D         15       17       DASS-D     <	14         14         BDI         Active           15         16         BDI         Passive           11         11         HADS-D         Passive           15         14         CES-DC         Active           10         12         BDI         Passive           6         8         CMDI         Passive           6         8         CMDI         Passive           14         17         BDI         Active           36         32         BDI         Active           49         44         CES-D         Passive           125         126         CES-D         Passive           19         11         RADS-2         Passive           28         29         HADS-D         Active           28         29         HADS-D         Active           49         50         BDI         Active           49         50         BDI         Active           49         50         BDI         Active           49         50         BDI         Passive           44         13         BDI         Passive           20 <t< td=""><td>14         14         BDI         Active         CBT           15         16         BDI         Passive         WL           11         11         HADS-D         Passive         WL           15         14         CES-DC         Active         MDT           10         12         BDI         Passive         WL           6         8         CMDI         Passive         WL           14         17         BDI         Active         CBT           36         32         BDI         Active         CBT           36         32         BDI         Active         PRT           49         44         CES-D         Passive         WL           125         126         CES-D         Passive         WL           19         11         RADS-2         Passive         WL           28         29         HADS-D         Active         AR           21         18         HADS-D         Active         TRT           49         50         BDI         Active         CBT           14         13         BDI         Passive         WL           45<!--</td--></td></t<>	14         14         BDI         Active         CBT           15         16         BDI         Passive         WL           11         11         HADS-D         Passive         WL           15         14         CES-DC         Active         MDT           10         12         BDI         Passive         WL           6         8         CMDI         Passive         WL           14         17         BDI         Active         CBT           36         32         BDI         Active         CBT           36         32         BDI         Active         PRT           49         44         CES-D         Passive         WL           125         126         CES-D         Passive         WL           19         11         RADS-2         Passive         WL           28         29         HADS-D         Active         AR           21         18         HADS-D         Active         TRT           49         50         BDI         Active         CBT           14         13         BDI         Passive         WL           45 </td

Note. Outcome measure abbreviations: BDI - Beck Depression Inventory, PAI-D - Personality Assessment Inventory Depression, HADS - Hospital Anxiety and Depression Scale, CES-CD-Centre for Epidemiological Studies Depression Scale for Children, CES-D-Centre for Epidemiological Studies Depression Scale, RADS-2 - Reynolds Adolescent Depression Scale 2, DASS – Depression Anxiety Stress Scale, PHQ9 – Patient Health Questionnaire 9 Control condition abbreviations: CT – Cognitive Therapy, IPP – Innovation Promotion Programme, ITSF – Intensive Twelve Steps Facilitation, WL – Waiting List, CBT – Cognitive Behavioural Therapy, MDT – Multidisciplinary Treatment Approach, PRT - Progressive Relaxation Training, AR - Applied Relaxation, TRT-, CBGT - Cognitive Behavioural Group Therapy, GDS-10=Geriatric Depression Scale; .

Table 4 Cumulative meta-analyses of ACT for Anxiety and Depression.

•	k	HOIS	At fina	l interim	analysis			Boundary crossed	Sufficiency	Additional sample required $(n)$
Primary Analysis	-		N	t	d	95% CI	$I^2$			
Anxiety/P	28	220	818	> 1	.95***	0.55-1.36	86.1	Y	Y	0
Anxiety/G	28	848	1628	> 1	.45*	0.19-0.64	84.1	N	Y	0
Depression/P	39	220	1037	> 1	.92***	0.64-1.19	82.7	N	Y	0
Depression/G	39	848	1987	> 1	.54***	0.34-0.73	80.6	Y	Y	0
Secondary Analysis										
Anxiety/P/Primary	5	220	146	.84	1.85**	0.05-364	93.3	N	N	74
Anxiety/G/Primary	5	848	275	.43	.77	-2.38 - 0.85	93.7	N	N	573
Anxiety/G/Active	10	848	558	.83	04	-0.21 - 0.14	5.3	N	N	290
Depression/P/Primary	12	220	360	> 1	1.22***	0.74-1.71	74.6	Y	Y	0
Depression/G/Primary	12	848	674	.71	.73**	0.30-1.16	63.9	Y	Y	0
Depression/G/Active	15	848	755	> 1	.26	-0.06 - 0.59	73.3	N	N	93

Note. /P - pre post comparison; /G - group comparison; /Primary - Primary target outcome; Active - Active control condition; k-number of studies; HOIS-Heterogeneity Adjusted Optimal Information Size; d-Pooled effect size; 95%CI-95% Confidence Interval; I<sup>2</sup>- I-squared value. As described above I<sup>2</sup> is defined as the ration between difference of Q and the degree of freedom by Q.

<sup>\*</sup> p < .05. \*\* p < .01. \*\*\* p < .001.

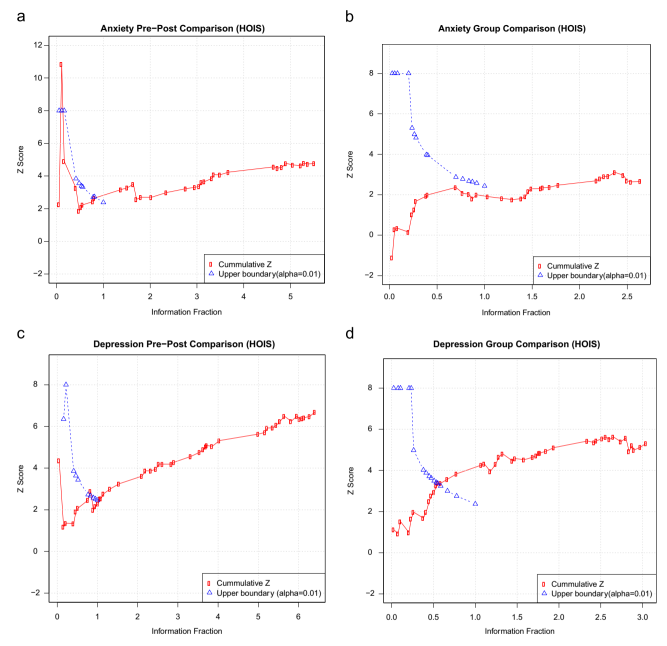


Fig. 2. Sequential meta-analyses of ACT for Anxiety and Depression (pre-post, group comparisons).

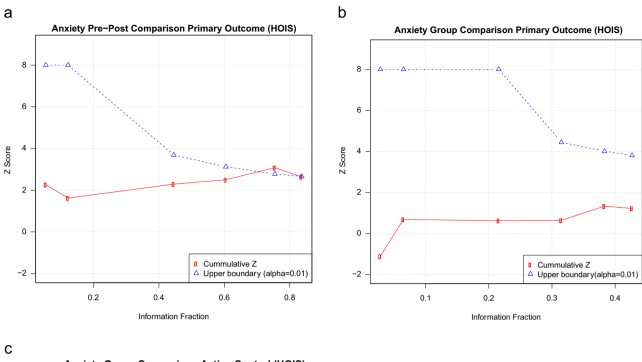
ranged from k=288 to k=671 studies, again suggesting absence of publication bias.

## 4. Discussion

The aim of this paper was to quantitatively review the cumulative evidence for ACT as a treatment for anxiety and depression. In doing so we generate sample size estimates for ACT trials in which there is currently insufficient evidence to determine the sufficiency of the evidence for ACT. Our novel statistical approach enables us to confidently appraise the treatment literature from a standpoint of statistical sufficiency. In contrast to conventional meta-analysis our approach controls for typical threats to statistical techniques in the evaluation of evidence based psychological therapies e.g. type I and II statistical errors; and between-sample

heterogeneity – thereby enhancing the statistical basis for determining treatment sufficiency.

In total we included k=28 and k=439 studies for anxiety and depression respectively. The cumulative pooled effect sizes for ACT for anxiety for both pre-post and group comparisons ranged from d=.45 to d=.95. In turn, the cumulative pooled effect sizes for depression in trials comparing ACT in pre-post and group comparisons ranged from d=.54 to d=.92. Our findings suggest that there is currently cumulative evidence for the efficacy of ACT versus controls in the treatment of anxiety and depression. All four cumulative meta-analyses for anxiety and depression were statistically significant. Equally findings from SMAs suggested sufficiency of evidence of an at least moderate effect for these conditions with the exception of ACT for anxiety in group comparisons. Thus one can conclude that with respect to the efficacy of ACT versus control conditions statistical sufficiency is reached and no further



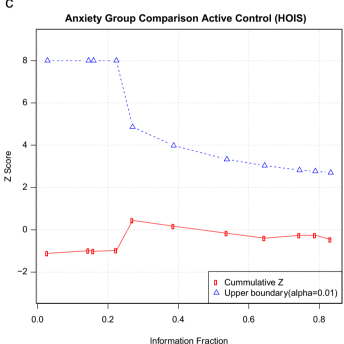


Fig. 3. Sequential meta-analyses of ACT for Anxiety (primary outcome, active control condition).

randomized clinical trials are required. Findings are thus in keeping with the previous literature for the overall cumulative efficacy of ACT (e.g. Hayes et al., 2006; Ost, 2008; Powers et al., 2009; Ruiz, 2010, 2012). However, applying an SMA approach suggests that although sufficiency has been reached in terms of clinical trials (as exemplified by above-threshold HOIS values), we conclude that the effect size of ACT for anxiety in-group comparisons is below a moderate effect.

However, findings are more qualified when considering the evidence for ACT compared to active control conditions i.e. existing evidence-based therapies, and when anxiety or depression were the primary treatment targets. For anxiety, although cumulative

evidence suggest a strong significant effect in pre-post comparisons ( $d\!=\!1.85,\,p\!<\!.001$ ) other cumulative meta-analyses for ACT for anxiety were not significant. Similarly, SMAs of ACT for anxiety for group comparisons failed to reach sufficiency (Fig. 3b and c). There is currently insufficient evidence to confidently infer a moderate effect between these intervention conditions (Fig. 3). However ACT for anxiety in pre-post comparisons, where ACT has been specified a-priori as the primary outcome, there is sufficient evidence to infer that ACT has a moderate effect. With respect to the analyses that fail to reach sufficiency two factors might account for these findings. Firstly, these SMAs for anxiety were underpowered; thus findings may have been erroneous. Secondly, Z values for the cumulative

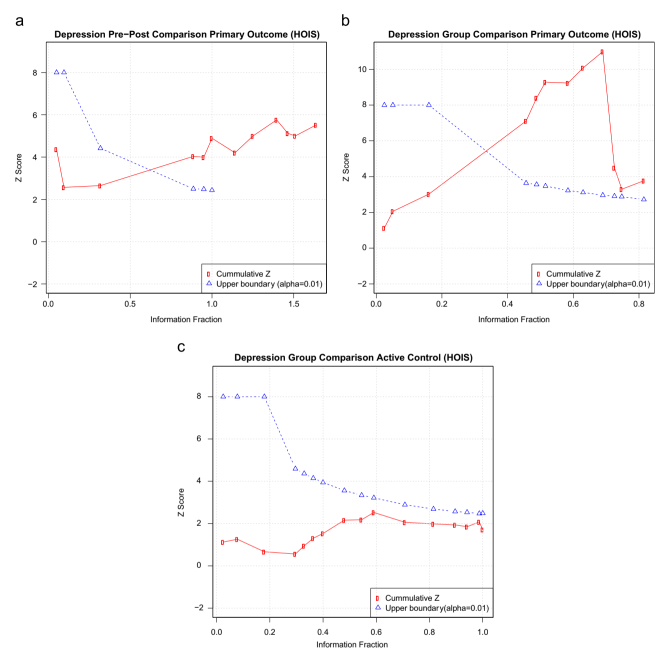


Fig. 4. Sequential meta-analysis of ACT for Depression (pre-post primary outcome, group primary outcome, group active control comparisons).

evidence suggest that, although HOIS has not yet been reached, there is no differential benefit for ACT in group conditions when ACT was predetermined as a treatment outcome or when compared to an active control ( $Z_{active} = -.46$ ; Fig. 3c). Equally, there is little differential benefit for ACT for anxiety in group comparisons when primarily targeting symptoms ( $Z_{group, primary} = 1.22$ ; Fig. 3b). Findings therefore seem to indicate similar outcomes are obtained irrespective of the active intervention. It is thus not likely to assume that additional clinical trials will reveal differential efficacy. These findings may inform treatment planning in settings or healthcare structures where estimates of differential efficacy are practically relevant to operationalizing treatment programmes.

Conversely, where ACT for depression was the primary outcome the cumulative evidence suggest large significant effects in both group-( $d_{group, primary}$ =.73, p < .01) and pre-post comparisons ( $d_{pre-post, primary}$ =1.22, p < .001). Findings from SMAs for these comparisons

also suggests that sufficient evidence exists to indicate an at least medium effect (Fig. 4a and b). However, cumulative evidence of ACT for depression against active controls suggests a small non significant effect ( $d_{group,\ active}$ =.26, n.s.). However. the SMA for this comparison (Fig. 4c) failed to reach statistical sufficiency. Consequently, the same issues raised above with regard to the anxiety comparisons that failed to reach sufficiency also apply to the effect of Act for depression when compared against normal controls.

It is noteworthy that careful consideration should be given whether and how additional randomized control trials meaningfully add to the existing knowledge base. For example in our study, when appraising the effect size of ACT for anxiety in active control conditions (i.e. d=-.04, n.s.), a difference in treatment gains might not become readily apparent. A possible explanation for this may be that ACT and CBT share therapeutic techniques, particularly exposure strategies.

Overall, the findings suggest that ACT is effective in the treatment of common mental health difficulties; however not more so than traditional treatment approaches. For instance, within the theoretical literature on ACT there has been discussion of the treatment mechanisms by which ACT leads to change in clinical presentations e.g. psychological flexibility. Our analysis does not evaluate these questions of process. Therefore, it may be a productive line of enquiry for future studies of ACT to emphasise the unique components of the intervention that may generate change in particular presentations (Arch and Craske, 2006). From this perspective, future research might thus attempt to answer questions of optimising treatment matching i.e. 'what works for whom' (Roth and Fonagy, 2006).

We acknowledge that the review is subject to several caveats. Efforts were made to retrieve grey literature in relation to ACT RCTs to minimise the impact of publication biases. Results of the analyses indicate no significant file drawer effect, thus publication errors are not likely to have significantly biased our results.

The heterogeneity indexes suggest a significant level of between study variance in the sample. Therefore it could be argued that the study sample is too diverse to meaningfully infer general conclusions regarding the evidence base as a significant proportion of between study variance has not been explained. However, our findings are in keeping with previous quantitative reviews and as such offer additional support for the veracity of our findings. In addition, the observed heterogeneity may be considered an accurate reflection of the scope of the application of ACT and as such might be suggestive of the external validity of our findings. We also acknowledge that our study set out to explore the efficacy of ACT for anxiety and depression. Consequently, the current analyses cannot comment on the efficacy of ACT for other complex mental health conditions e.g. psychosis (White, 2013) or physical health conditions such as chronic pain (McCracken et al., 2013).

The use of SMA has introduced discussion around the use stop criteria in research considerations (Higgins et al., 2010; Pogue and Yusuf, 1997). As noted our intention in using the method has been to inform future research. Although historically SMA has been used to inform decision on research funding in clinical trials (Wetterslev et al., 2008) we caution against the indiscriminate application of SMA for such sole purpose. Whether or not individual trials proceed is a complex decision making process of which sufficiency considerations might comprise one aspect of multiple evidence strands, e.g. in conjunction with Disability Adjusted Life Years (DALYs).

Lastly, it is important to note that symptom reduction is not the primary intervention objective of ACT interventions; instead ACT aims to improve psychological flexibility (Harris and Hayes, 2009; Hayes et al., 2011). Future studies might want to investigate other outcome domains consistent such as function or quality of life which may be more germane to third wave interventions.

#### **Conflict of interest**

The authors report no conflicts of interest in the preparation of this article. Dr MacBeth was supported by an NHS Research Scotland Career Research Fellowship while preparing the article. The funder was not directly involved in the conduct of the research or the write-up.

## Acknowledgement

Kuppens et al. (2012) whose previous publications introduced us to SMA and thus enabled us to use the method for the purpose of this meta-analysis.

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